WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 4:		(11) International Publication Number:	WO 89/ 00606
C12P 7/64, C09F 5/02 A61K 7/00	A1	(43) International Publication Date: 26 January	nary 1989 (26.01.89)

(21) International Application Number	r: PCT/US88/02483	(72) Inventor; and
(22) International Filing Date:	20 July 1988 (20.07.88)	(75) Inventor/Applicant (for US only): LONG, Thomas, Veach, II [US/US]; 315-B Summer Rest Road, Wil-

mington, NC 28403 (US). (31) Priority Application Number: 075,662 (74) Agent: BARBER, Lynn, E.; Olive & Olive, P.O. Box 2049, Durham, NC 27702 (US). 20 July 1987 (20.07.87)

(81) Designated States: AT (European patent), AU, BB, BE (European patent), BG, BJ (OAPI patent), BR, CF (OAPI patent), CG (OAPI patent), CH (European patent), CM (OAPI patent), DE (European patent), DK, FI, FR (European patent), GA (OAPI patent), GB (European patent), HU, IT (European patent), JP, KP, KR, LK, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL (European patent), NO, RO, SD, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent), US. (33) Priority Country: (60) Parent Application or Grant (63) Related by Continuation 075,662 (CIP) Filed on 20 July 1987 (20.07.87) (71) Applicant (for all designated States except US): MARI-CULTURA, INCORPORATED [US/US]; Post Office Drawer 565, Wrightsville Beach, NC 28480 (US). patent), US.

Published

With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: MICROORGANISM PRODUCTION OF OMEGA-3 (N-3) LIPIDS

(57) Abstract

(32) Priority Date:

The invention comprises the use of obligately and facultatively marine eukaryotic microorganisms for the production of Omega-3 (n-3) fatty acids that may be used in food, cosmetic, and pharmaceutical products. In the invention the microorganisms are grown heterotrophically, harvested, and extracted for lipid products.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

	. •				
AT	Austria	FR	France	ML	Mali
UΑ	Australia	GA	Gabon	MR	Mauritania
BB	Barbados	GB	United Kingdom	MW	Malawi
BE	Belgium	HU	Hungary	NL	Netherlands
BG	Bulgaria	IT	Italy	NO	Norway
BĴ	Benin	JP	Japan	RO	Romania
BR	Brazil	KP	Democratic People's Republic	SD	Sudan
CF	Central African Republic		of Korea		Sweden
CG	Congo	KR	Republic of Korea	SN	Senegal
CH	Switzerland	LI	Liechtenstein	SU	Soviet Union
CM	Cameroon	LK	Sri Lanka	TD	Chad
DE	Germany, Federal Republic of	LŪ	Luxembourg	ΤĞ	Togo
DK	Denmark	MC	Monaco	US	United States of America
FI	Finland	MG	Madagascar	00	Ourred prates of Willettes

15

20

25

30

MICROORGANISM PRODUCTION OF OMEGA-3 (N-3) LIPIDS

Field of the Invention

This invention relates to fatty acid production from microorganisms. In particular, this invention relates to the use of obligately or facultatively marine eukaryotic microorganisms grown heterotrophically and the production of a group of highly polyunsaturated fatty acids known as Omega+3 or n-3 fatty acids.

Background Information

The biosynthesis of fats and oils by microbes such as yeasts, bacteria, molds, and algae is well established, and inventors have devised ways of optimizing the growth conditions for this biosynthesis. For example, yeasts or molds in mixed culture with bacteria and grown on mixtures of carbohydrates or hydrocarbons under aerobic fermentation conditions were found to produce numerous amino acids plus, in some cases, unspecified oil and fat. (U.S. Pat. No. 3,793,153). The disclosure of this reference and all others cited herein are hereby incorporated by reference. such as Candida and Rhodotorula also produce single-cell protein and undifferentiated lipid from vegetable carbohydrates including starch (U.S. Pat. No. 4,230,806). Under aerobic conditions Candida tropicalis produces unsaturated dicarboxylic acids having 14 to 22 carbon atoms from unsaturated fatty acids or their esters (U.S. Pat. No. 4,474,882). Numerous yeasts dramatically increase their production of 1, 3-disaturated-2-unsaturated triglycerides, a predominant component of cacao butter, under high oxygen fermentation conditions in a growth medium containing one or more fatty acids having between 10 and 20 carbon atoms. Pat. No. 4,485,173). Rhodococcus rhodochrous (formerly

10.

15

30

35

Nocardia aurantia) synthesizes hydroxy-fatty acids and keto-fatty acids, such as 10-hydroxy-12-octadecenoic acid and 10-hydroxysteric acid, under aerobic conditions by hydration of unsaturated fatty acids such as linoleic and oleic acid, respectively. (U.S. Pat. No. 4,582,804). Methods have also been devised to increase oil production and recovery from halophilic algae such as Dunaliella by growing the cells photosynthetically in a saline solution with a growth promoting enzyme and using water insoluble solvent extraction. The oils obtained had a carbon and nitrogen content similar to crude oil and were apparently saturated fatty acids and wax esters. (U.S. Pat. No. 4,341,038).

Although the inventors of the cited patents and other investigators have determined ways to obtain and optimize fat and oil production from certain microorganisms, none of these methods produces polyunsaturated fatty acids, and, in particular, they do not yield the highly polyunsaturated fatty acids designated Omega-3 or n-3 fatty acids. The Omega-3 and Omega-6 fatty acids are the two principal classes of polyunsaturated fatty acids. The Omega=3 (or n=3) fatty acids have the first unsaturation (double bond) at the third carbon from the methyl (or Omega) end of the fatty acid, and the Omega=6 (or n=6) fatty acids have the first unsaturation at the sixth carbon from the methyl end of the molecule. Omega=6 fatty acids, such as linoleic acid, are common components of vegetable oils, such as corn oil. The microbial production of linoleic acid by culturing fungi of the Pellicularia genus has been elaborated (U.S. Pat. No. 4,281,064).

The Omega-3 fatty acids are prominent components of some fish oils, although they are not synthesized by the fish but, instead, appear to be absorbed from their diets. These highly polyunsaturated fatty acids have been shown effective in humans in preventing or remedying cardiovascular disease and a number of auto-immune diseases, such as arthritis. They

ń

5

10

15

20

25

30

appear to be an essential nutrient. Although an earlier fermentation process of the prokaryotic Streptomyces organisms (U.S. Pat. No. 3,127,315) produced a hypocholesterolemic agent referred to as M=850, it is clear from the extraction procedures used that this agent is not related to the Omega-3 fatty acids or their action.

The focus of medical research on the fatty acids has been principally on two of the Omega-3 fatty acids, eicosapentaenoic acid (EPA: a 20-carbon fatty acid having 5 unsaturations) and docosahexaenoic acid (DHA: a 22-carbon fatty acid having 6 unsaturations), although others of this class may prove to be important. It is widely recognized that the principal dietary sources of these chemical moieties for fish are photosynthetic algae and microalgae. The use of the marine microalga, Chlorella minutissima, to produce Omega-3 fatty acids photoautotrophically has been the subject of recent patents (U.S. Pat. No. 4,615,839 and Jap. Pat. Discl. No. Sho-61-63624). A process for preparation of eicosapentaenoic acid from linolenic acid using enzymes from microalgae and macroalgae also has been described (Jap. Pat. Discl. No. Sho-61-31092).

In contrast, no method has been described heretofore to produce Omega-3 fatty acids from microbes that are grown heterotrophically, using as a nutrient a sugar, carbohydrate, or other source of "pre-formed" carbon. Although a process for cultivating the freshwater microalga Chlorella mixotrophically on lower fatty acids to aid in disposing of organic wastes (U.S. Pat. No. 3,444,647) was developed, no product description is provided, and this species does not produce Omega-3 fatty acids.

Summary of Invention

This invention comprises use of certain heterotrophic and autotrophic eukaryotes grown heterotrophically to produce

25

lipid that contains Omega-3 fatty acids as prominent constituents, and in use of this lipid or its constitutents in nutritional supplements; in foods for humans and other animals, including aquacultured species; and in drugs and pharmaceuticals. No prokaryote is known to synthesize Omega-3 fatty acids. Eukaryotic synthesis based on autotrophic growth of photosynthetic microalgae presents problems of light and nutrient provision that impair economic production. inventor has determined that obligately or facultatively marine eukaryotes, including but not limited to fungi, thraustochytrids, microalgae (in particular, diatoms and dinoflagellates), and yeasts produce high concentrations of docosahexaenoic acid and eicosapentaenoic acid when grown under heterotrophic growth conditions. Obligately freshwater (aquatic) heterotrophic eukaryotes do not appear to produce these chemicals.

Conventional sources of Omega-3 fatty acids are oils from such fish as salmon, anchovies, sardines and menhaden. These fish oils in the unrefined form contain cholesterol. Depending on the environment and diet of the fish, these fish oils may also contain heavy metals and synthetic organic chemicals such as polychlorinated biphenyls (PCB's), polybrominated biphenyls (PBB's), dieldrin and aldrin. In addition to these disadvantages, the fish oils have an unpleasant fishy odor and taste. It is a further advantage of this invention to provide lipids containing Omega-3 fatty acids that are not contaminated with heavy metals or undesirable synthetic organic chemicals. The invention also provides Omega-3 fatty acid products that do not have a strong fishy odor and taste.

Detailed Description of the Invention

This invention represents that selected heterotrophic eukaryotes that are either halo-tolerant or halophilic, when cultured, cultivated, or fermented in a salt-containing medium

10

15

20.

25

30

35

or one containing natural or artificial seawater, will produce Omega=3 fatty acids, including but not limited to EPA and DHA, as a significant percentage of total lipid. Furthermore, the use of such microorganisms yields a lipid fraction that is useful as a nutritional supplement; as a food additive in margarines, cooking oils, salad dressings, baked products, infant nutritional formulae and adult enteral nutritional formulae; as a skin care product or cosmetic; as a drug or pharmaceutical; as a component of an intravenous, parenterally administered fluid; and as an animal or aquaculture feed or feed additive. The high levels of Omega-3 fatty acids produced by these marine eukaryotes grown heterotrophically are unique. Prokaryotic microorganisms generally do not produce these fatty acids. Although photosynthetic, autotrophic eukaryotes do synthesize the Omega-3 fatty acids, the rates of growth of these organisms and their production of the fatty acids under photosynthetic growth conditions are significantly less than when heterotrophic eukaryotes are cultivated heterotrophically. Specific heterotrophs appropriate for use in this invention include, but are not limited to: the thraustochytrids, Thraustochytrium roseum and T. aureum; the phycomycetes fungi, Pythium sp. and Schizochytrium aggregatum; the diatom, Nitzschia sp.; and the dinoflagellate, Crypthecodinium cohni. These species are maintained in the American Type Culture Collection and the algae culture collection of the University of Texas at Austin.

The Omega-3 fatty acids appear to be produced only by marine microorganisms or by halophilic or halo-tolerant species. Thus, in the preferred embodiment, either seawater, an artificial seawater, or other saline solution is used as the solvent in the culture medium. The complete medium is referred to as a saline culture medium. The carbon source in this saline culture medium may be a relatively simple carbohydrate source, for example, glucose, sucrose, mannose, or molasses, or, if slower cultivation is permitted, vegetable fibers such as grasses or bagasse.

30

After inoculation of the saline culture medium with the selected eukaryotic microorganisms, the medium is incubated under conditions favorable for heterotrophic growth of the microorganism. Extractions of the culture are accomplished by solvent extraction using a mixed organic solvent and standard techniques. This invention will be more clearly understood by references to the following illustrative examples, which are not to be construed as limiting the invention:

EXAMPLE 1

Pure cultures of the thraustochytrids, yeasts, fungi, or microalgae are inoculated into liquid media of successively larger volumes starting at 100 ml and staging up to larger cultures. A saline culture medium is prepared by mixing 1.0 g of glucose with 0.1 g yeast extract in 1000 ml of aged seawater. The amount of glucose can be increased to 5 g and the pH in the preferred embodiment is adjusted to between 7 and 7.5. The cultures are grown for two weeks at approximately 25 to 28° C. to facilitate heterotrophic growth. Growth vessels are either shaken with rotary shakers or magnetically stirred. The species may be grown either unilluminated or illuminated with moderate light, even though the organisms are growing heterotrophically. Harvesting is carried out by centrifugation or freeze drying.

These microorgansims may be extracted in the wet state directly after harvesting or in the freeze dried state, using a mixture of non-polar and polar organic solvents consisting of methanol, chloroform, and water in the proportions 2:1:0.8. The solvents are mixed with the microorganisms and allowed to stand for 1/2 to 3 hours. After this period, additional chloroform and water are added to yield a solvent ratio of 2:2:1.8 of methanol, chloroform, and water. The chloroform layer contains the total lipid fraction, which is comprised of a sufficiently high concentration of Omega-3 fatty acids to be useful in nutrition and medicine.

20

25

30

Example 2

The procedures followed in Example 1 are used, except that the saline culture medium is composed of 1 to 5 g glucose, 1 g yeast extract, and 1 g peptone in 1 liter of seawater. The pH is adjusted to between 7 and 7.5.

Example 3

The procedures followed in Example 1 are utilized, except that an artificial seawater base is prepared for this saline culture medium. This consists of 2.5 g NaCl, 0.5 g MgSO4. 7H2O, 0.1 g KCl, 0.01 g KH2PO4, 0.02 g CaCO3 and sufficient H2SO4 to dissolve the above compounds. To this solution, (NH4)2 SO4 is added in the amount of 0.02 g, along with 0.2 with 0.2 g NaH-glutamate, 0.1 g Agar, 1.0 ug Thiamine-HCl, 0.1 ug Cyanocobalamin, 5.0 mg Na2EDTA, and the trace metals 0.05 mg FeSO4.7H2O, 0.02 mg ZnSO4.7H2O, 0.01 mg MnSO4.H2O, 2.0 ug CoSO4.7H2O, 0.2 ug CuSO4.5H2O, 2.0 ug H3BO3, 2.0 ug NaMoO4.H2O with sufficient distilled water to yield 100 ml of solution. To this is added 0.1 to 0.5 g glucose or other sugar, 0.01 g NaHCO3, and the pH is adjusted to between 7 and 7.5.

Example 4

The media and procedures employed in any of the above examples are used except that the cultures are exposed to light of moderate intensity. Heterotrophic growth of certain marine eukaryotes, such as the thraustochytrids, is enhanced under such conditions.

Example 5

The media and procedures employed in any of the above examples are used except that additional limited quantities of available N and P (such as 0.085 g NaNO3 and 0.012 g NaH $_2$ PO $_4$) are added to the saline culture medium. Heterotrophic growth

of certain marine eukaryotes, such as the microalgae, is enhanced in this medium.

The samples harvested from these examples produce lipid fractions containing Omega-3 fatty acids. After extraction and esterification to form the methyl esters, gas chromatographic analyses show that the Omega-3 fatty acids may constitute as much as 10 to 50% of the total fatty acid fraction. They are generally contained in phospholipids, glycolipids, mono-, di-, or triglycerides, and sulfolipids, or as the free acids, but are not limited to these forms.

¥

15.

20

What is claimed is:

- A process of using heterotrophically grown obligately or facultatively marine eukaryotic microorganisms as a source for the production of Omega=3 (n=3) fatty acids.
- 2. A process according to claim 1, wherein the marine eukaryotic microorganism is selected from the group consisting of thraustochytrids, lower fungi, yeasts, and microalgae.
- 3. A process for the production of Omega=3 (n=3) fatty acid products from obligately or facultatively marine eukaryotes, comprising the steps of:
 - (a) inoculating a saline culture medium containing a carbon source with marine eukaryotic microorgansims;
 - (b) incubating the inoculated saline culture medium under conditions conducive to heterotrophic growth of the marine eukaryotic microorganisms;
 - (c) harvesting the marine eukaryotic microorganisms from the saline culture medium; and
 - (d) extracting the lipid fraction from the harvested marine eukaryotic microorganisms.
 - 4. A process according the claim 3, wherein the saline culture medium comprises seawater.
 - 5. A process according to claim 4, wherein the carbon source comprises glucose.
 - 6. A process according to claim 4, wherein the conditions conducive to heterotrophic growth comprise a pH of from 7

- to 7.5 and a temperature of 25 to 28° C.
- 7. A process according to claim 3, wherein the saline culture medium comprises an artificial seawater base.
- 8. A process according to claim 7, wherein the carbon source comprises glucose.
 - 9. A process according to claim 7, wherein the conditions conducive to heterotrophic growth comprise a pH of from 7 to 7.5 and a temperature of 25 to 28° C.
- 10. A process according to claim 3, wherein the conditions conducive to heterotrophic growth comprise exposure to light of moderate intensity.
 - 11. A fatty acid product comprising an Omega-3 (n-3) fatty acid extracted from a heterotrophically grown, obligately or facultatively marine, eukaryotic microorganism culture.
 - 12. The fatty acid product of claim 11, wherein the fatty acid product may be used as a nutritional additive to the diet of humans.
- 13. The fatty acid product of claim 11, wherein the fatty acid product may be used as a pharmaceutical product.
 - 14. The fatty acid product of claim 11, wherein the fatty acid product may be used as a skin care or cosmetic product.
- 15. The fatty acid product of claim 11, wherein the fatty
 25 acid product may be used as an animal feed additive.

INTERNATIONAL SEARCH REPORT

International Application No. PCT/US88/02483

I. CLASS	SIFICATIO	N OF SUBJECT MATTER (if several classi	fication symbols apply, indicate all) ⁶	
According	to Internat	Ional Patent Classification (IPC) or to both Nat	ional Classification and IPC	
IPC(4	4): C1	2P 7/ 64; C09F 5/02;	ADIK //UU	
		435/134; 260/405.5;	420/60	
II. FIELD	S SEARCE			
		Minimum Docume		
Classificati	on System	· · · · · · · · · · · · · · · · · · ·	Classification Symbols	
U.S.		435/134, 254, 257, 426/60; 260/405.5;	911, 946, 171, 255 514/844, 772	
		Documentation Searched other to the Extent that such Documents	han Minimum Documentation are included in the Fields Searched ⁸	
1988;	; File	Chemical Abstracts S Biosis 1969-1988 US ment for search terms	Patent File (USPAT)	e CA, 1967-
		ONSIDERED TO BE RELEVANT 9		
Category *		ion of Document, 11 with indication, where app	ropriate, of the relevant passages 12	Relevant to Claim No. 13
х	Ass	ournal of The Canadian sociation, (Toronto, C plume 143, issued Apri	anada),	12 and 13-15
	(ACKMAN, R.G.), "Polyunsaturated Fatty Acid Content of Kelp Tablets and Dulse", pages 150-154. See entire document.			
X Y	Lancet (London, England), Volume 2, 1ssued 15 July 1978, (DYERBERG ET AL). "Eicosopentaenoic Acid and Prevention of Thrombosis and Atherosclerosis?," pages 117-119. See entire document.			
			•	
	· .			
·	!			
*Special categories of cited documents: 10 "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention				
filin "L" doc whi	g date ument whic ch is cited	h may throw doubts on priority claim(s) or to establish the publication date of another r special reason (as specified)	cannot be considered novel or involve an inventive step "Y" document of particular relevance cannot be considered to involve a	cannot be considered to e; the claimed invention an inventive step when the
"O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed "A document member of the same patent family				
IV. CERTIFICATION				
Date of the	Actual Co	mpletion of the International Search	Date of Mailing of this International Sec	arch Report
30 Au	gust	1988	.1 2110. 100	
b .	al Searchin	g Authority	Signature of Authorized Officer	,
TSA/11	21	•	IRENE MARX	

	ENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEE	Relevant to Claim No
alegory •	Citation of Document, with indication, where appropriate, of the relevant passages	Transfer to Gram 140
x	Bulletin of the Japanese Society of	15
$\frac{\mathbf{X}}{\mathbf{Y}}$	Scientific Fisheries (Tokyo,	12-13
*	Japan), Volume 46, issued	
-	1980, (WATANABE ET AL.)	
	"Relationship between Dietary	·
	Value of Brine Shrimp Artemia salina	
ŀ	and their content of omega-3.	
	Highly unsaturated Fatty acids",	
ļ.	pages 35-41. See entire	
1	document.	
Y	US, A, 4,661,343 (ZABOTTO)	14
Y		1 **
	28 April 1987, See columns	·
-	4 and 5 in particular.	ļ
	Towns 1 of Dyshamoslogic (Athona	1-3, and
$\frac{\mathbf{X}}{\mathbf{Y}}$	Journal of Protozoology (Athens,	5-8
Y	Georgia), Volume 17, issued	
* ** .	1970 (HARRINGTON ET AL.) "The	4-6,9
	Polyunsaturated Fatty Acids	and 10
	of Marine Dinoflagellates",	•
	pages 213-219. See entire	\. ·
	document.	
X	Comparative Biochemistry and	1-3, 7
Ÿ	Physiology, (Elmsford, New York),	and 10
ŀ	Volume 29, issued 1969,	4-6
	(ELLENBOGEN, B. ET AL,)	and
- -	"Polyunsaturated Fatty	8-9
	Acids of aquatic fungi: possible	• •
-	phylogenetic significance". pages	
	805-811. See entire document.	
ì		
$\frac{\mathbf{X}}{\mathbf{Y}}$	Journal of Experimental	1-3,7
Y	Botany (Oxford, England)	and 10
	Volume 25, issued August 1974	4-6
-1	(OPUTE, F.I.), "Lipid and	and
	Fatty Acid Composition of Diatoms"	8-9
	pages 823-835. See entire	·
. [document.	
, İ		
Y	Mycologia (Bronx, New York), Volume 55,	1-10
<u> </u>	issued 1963 (GOLDSTEIN, S.)	
" .	"Studies of a new species of	
. 1	Thraustochytrium that displays	· .
.	light stimulated growth",	1
	pages 799-811. See entire document.	•
- 1		
j		
1		

Search Terms

Nitzschia

Elcosapentaenoic docosahexaenoic cosmetic or skin care polyunsaturated or unsaturated animal (feed or rood) (omega-3 or n-3) fatty acid lipıd halotolenant or halophilic marine, salin?, salt fungus, fungi, alga? microbe, microorganism diatom, dinoflagellate ferment heterotroph thraustochytrid phycomycetes pythlum schizochytrium